Bimodal Distribution of Erythrocytes in Heterozygotes for Strong Mediterranean Glucose-6-phosphate Dehydrogenase Deficiency*

E. SARTORI, F. PANIZON, and F. ZACCHELLO

From the Pediatric Clinic, University of Pavia, Pavia, Italy

Haemolytic favism is a condition that affects children predominantly, and is frequent in Sardinia and in other Mediterranean areas. It is related to a deficiency of the enzyme glucose-6-phosphate dehydrogenase (G6PD) in the erythrocytes, as is 'primaqueine sensitivity' of American Negroes (Larizza, Brunetti, Grignani, and Ventura, 1958; Siniscalco, Motulsky, Latte, and Bernini, 1960; Carson, Flanagan, Ickes, and Alving, 1956; Gross, Hurwitz, and Marks, 1958; Tarlov, Brewer, Carson, and Alving, 1962). In both conditions the deficiency is due to a mutant allele of an X-linked gene, but in haemolytic favism the enzyme is of an electrophoretically slow (B) type, while in primaquine sensitivity it is of a fast (A) type. The deficiency is definitely more pronounced in haemolytic favism, but recently a mild deficiency of the B type has been found in Greece (Kirkman, 1962; Kirkman, Schettini, and Pickard, 1964; Porter, Boyer, Watson-Williams, Adam, Szeinberg, and Siniscalco, 1964; Stamatoyannopoulos, Panayotopoulos, and Papayannopoulou, 1964).

In spite of the different dose of X-linked alleles, hemizygous males and homozygous females, both normal and mutant, do not differ in their mean G6PD enzyme levels (Adinolfi, Bernini, Carcassi, Latte, Motulsky, and Siniscalco, 1960; Davidson, Childs, and Siniscalco, 1964) nor are there dosage differences among individuals with abnormal numbers of sex chromosomes, such as XO, XXY, XXX, XXXY, XXXX (Grumbach, Marks, and Morishima, 1962; Harris, Hopkinson, Spencer, Court Brown, and Mantle, 1963).

In both the American and Mediterranean deficiencies, heterozygous females present enzyme levels that are intermediate between but also overlap with those of normal and of mutant hemizygous males (Beutler, Yeh, and Fairbanks, 1962; Adinolfi, Davidson, Latte, Meera Khan, Piomelli, Ratazzi, and Siniscalco, 1963; Davidson et al., 1964). Moreover, Sardinian females clinically affected with haemolytic favism are observed with a frequency that is higher than the number of homozygotes and lower than the number of heterozygotes expected on the basis of the local gene frequencies (E. Sartori, unpublished data).

These discrepancies could be explained by the Lyon hypothesis (1962) which postulates that one of the two X chromosomes of normal females is genetically inactive and that either of the two X chromosomes may be inactivated in early embryonic life at random in different cells of the same individual.

According to Beutler et al. (1962) an implication of the hypothesis is the existence, in the G6PD deficient heterozygotes, of two distinct cell populations, one normal and one deficient. This assumption has been submitted to experimental tests by many authors. Among those who confirm the hypothesis, some have not considered sufficiently the possibility that the erythrocytes can behave as if they were composed of two distinct populations also in hemizygous males, provided that their behavior is referred to an appropriate single measurement. Thus it was well known that in primaquine-sensitivity the haemolysis was clinically and haematologically self-limited and that the limit was related to cell age as well as to the dose of the drug (Dern, Beutler, and Alving, 1954; Kellermeyer, Tarlov, Schrier, Carson, and Alving, 1961). The haemolysis is self-limited also in haemolytic favism (Sartori and Panizon, 1957). Furthermore Frischer, Bowman, and Carson (1964) and Papayannopoulou and Stamatoyannopoulos (1964), applying the cyanmethaemoglobin

Received May 20, 1965

* This research has been supported in part by C.N.R.
Erythrocytes in Heterozygotes for G6PD Deficiency

elution technique to the methaemoglobin reduction test of Brewer, Tarlov, and Alving (1960), were able to visualize microscopically two kinds of erythrocytes in mild G6PD-deficient males as well as in intermediate females, as found by Sansone, Rasore-Quartino, and Veneziano (1963) and by Tonz and Rossi (1964).

These objections do not apply to the experiments of Beutler et al. (1962) who reported that the disappearance of reduced glutathione from the erythrocytes of females with intermediate red-cell G6PD activity challenged with acetylphenylhydrazine and the reduction of their methaemoglobin occurred at two different rates, as in artificial mixtures of enzyme-deficient and normal erythrocytes. In our hands the methaemoglobin reduction as used by these authors failed to give consistent results. More recently, Beutler and Baluda (1964), using blood of two Negro females with intermediate red-cell G6PD-activity and with the sickle-cell trait, succeeded by means of sickling and successive millipore filtering in separating the erythrocytes in which methaemoglobin was still present from those in which it had been reduced. They found that the filtered not sickled methaemoglobin-rich erythrocytes were present in relevant amount and had a very little G6PD-activity. Using another approach, Davidson, Nitowsky, and Chilids (1963) had previously demonstrated two kinds of cells by cloning skin biopsies of Sardinian women heterozygous for quantitative enzyme deficiency and of American Negro women heterozygous for qualitative electrophoretic enzyme variants.

Evidence against the hypothesis derives in the first place from failure to detect in the blood of heterozygotes a portion of DF 32 P-labelled erythrocytes behaving in vivo as normal ones (Brewer, Tarlov, and Powell, 1962), and secondly for failure to separate from the blood of double heterozygotes Xg (a+) and Xg (a−) erythrocytes with different G6PD activities, which could easily be separated from artificial mixtures of hemizygous bloods (Gorman, Dire, Treacy, and Cahen, 1963).

As to this negative evidence, we think that the long-term survival behaviour in vivo of a double population of erythrocytes can hardly be predicted with sufficient precision. Furthermore, the Xg 0 locus may belong to a non-inactivated region of the X chromosome. Such an hypothesis has been formulated by Russell (1963) to explain some variegated type position effects in the mouse, by limitation in spread and in region of origin of the inactivation of the mammalian X chromosome.

In conclusion, the existence of a true erythrocyte mosaic in females heterozygous for G6PD-deficiency is still open to some doubt, and further investigations are desirable.

Given the appropriate variable, two normal distributions of it found in heterozygotes would prove the mosaic, while a single distribution would exclude it. We searched accordingly for a variable that would give normal distributions for both normal and mutant hemizygous erythrocytes and first tried with primaquine. Panizon and Zacchello (1965) in fact observed that erythrocytes of enzyme-deficient males, labelled with 51Cr and then transfused, underwent an almost complete destruction within one day when first incubated with 0.10 mg./ml. primaquine, while only about half the deficient cells were destroyed if the drug was lowered to 0.02 mg./ml. Primaquine, however, proved to be unsuitable, since at higher doses it modifies the blood physically, and it was, therefore, impossible to obtain a distribution for normal cells. We finally succeeded in our purpose by first damaging the erythrocytes in vitro with increasing concentrations of menadione sodium bisulphite, the hydrosoluble vitamin K 3, and then destroying the damaged ones in vitro by means of transfusing the blood into compatible recipients, after having labelled it with 51Cr. The proportion of destroyed cells was finally inferred from short-term survival determinations.

Material and Methods

Subjects. Blood was drawn from 8 healthy Italian adults, Wassermann negative and without history of malaria. Four were males, two of them normal and two strongly deficient for erythrocyte G6PD-activity. The other four were females heterozygous for the mutant gene. In three, heterozygosity was proven because each had at least one normal and one enzyme-deficient son; in the fourth, heterozygosity was strongly presumptive because her father had suffered from haemolytic favism, but she presented only minor fava-induced disorders and repeatedly showed intermediate values for erythrocyte G6PD activity. Recipients were normal volunteers, compatible for ABO and Rh blood groups.

Glutathione and Enzyme Determinations. Erythrocyte-reduced glutathione(GSH), before and after incubation (for 2 hours at 37° C. with 5 mg./ml. acetylphenylhydrazine and 2 mg./ml. glucose), was determined by the method of Beutler, Duron, and Kelly (1963). Studies were carried out in the Bausch and Lomb spectrophotometer and expressed as mg./100 ml. packed erythrocytes.

Erythrocyte G6PD activity was measured with the 'Boehringer Test Combination', using the Beckman DB spectrophotometer with recorder, at a temperature of 18–22° C. The activity was expressed as optical density units per minute/100 g. Hb. Erythrocyte acid phosphomonoesterase (APM) was determined by the method of Oski, Shahidi, and Diamond (1963) and expressed as mg. phenol in 30 minutes per 100 g. Hb. Erythrocyte catalase activity was tested by the perborate method of Feinstein as used by Tarlov and Kellermeyer (1961) and expressed as mEq of NaBO 4 decomposed during the first minute.

The results of all these determinations in the blood of the eight donors are listed in Table 1.
TABLE I
GLUTATHIONE AND ENZYME DETERMINATIONS IN 2 NORMAL, 2 G6PD-DEFICIENT, AND 4 HETEROZYGOUS FEMALES

<table>
<thead>
<tr>
<th>Subject No., Sex, and Age (y.c.)</th>
<th>GSH</th>
<th>G6PD</th>
<th>APM</th>
<th>Catalase</th>
<th>MSB*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 M 25f</td>
<td>61</td>
<td>86</td>
<td>41</td>
<td>66</td>
<td>81</td>
</tr>
<tr>
<td>2 M 26f</td>
<td>65</td>
<td>86</td>
<td>41</td>
<td>66</td>
<td>81</td>
</tr>
<tr>
<td>3 M 26f</td>
<td>35</td>
<td>1070</td>
<td>0</td>
<td>710</td>
<td>81</td>
</tr>
<tr>
<td>4 M 26f</td>
<td>41</td>
<td>0</td>
<td>967</td>
<td>710</td>
<td>81</td>
</tr>
<tr>
<td>5 F 21f</td>
<td>66</td>
<td>710</td>
<td>967</td>
<td>710</td>
<td>81</td>
</tr>
<tr>
<td>6 F 28f</td>
<td>51</td>
<td>710</td>
<td>967</td>
<td>710</td>
<td>81</td>
</tr>
<tr>
<td>7 F 26f</td>
<td>22</td>
<td>710</td>
<td>967</td>
<td>710</td>
<td>81</td>
</tr>
<tr>
<td>8 F 21f</td>
<td>81</td>
<td>710</td>
<td>967</td>
<td>710</td>
<td>81</td>
</tr>
</tbody>
</table>

* 24-hour residual percentage of 14Cr-labelled erythrocytes transfused after incubation with 1 × 10^-4 M menadione sodium bisulphite.
† G6PD deficient.
‡ Heterozygous.

Incubation and Labelling. Standards having been derived from preliminary experiments, 32 mg. menadione sodium bisulphite (MSB) were dissolved each time in 10 ml. phosphate-buffered saline at pH 7.40 and sterilized by means of boiling. This solution was brought from 320 down to 1-25 mg./100 ml. by serial twofold dilutions. To 5 ml. blood was added 1 ml. solution and the following range of final concentrations, expressed in Moles × 10^-4, was obtained: 1; 1/2; 1/4; 1/8; 1/16. From each donor, 7 samples of heparinized blood, fresh or stored for no longer than 12 hours at about 4°C, to which were added 1% of a sterile 20% glucose solution, were tried against consecutive dilutions of MSB, ranging from 1 to 4 × 10^-4 M for the G6PD-deficient, from 1 to 8 × 10^-4 M for the intermediate, and from 1 to 16 × 10^-4 M for the G6PD-normal donors (see Table II). For all samples the incubation time was 3 hours. At the third hour 1 μC 14Cr salt/kg. of the recipient’s body weight was added to every specimen, which was then incubated for a further 30 minutes. Finally the plasma was removed by gentle centrifugation and the erythrocytes were suspended in sterile saline and injected intravenously.

Recording. For each donor seven recipients were injected, and from these blood was drawn in duplicate 10 minutes and 24 hours after the injection. The radioactivity of the specimen was then measured simultaneously for all those belonging to the same donor. The mean value after 24 hours was referred to that after 10 minutes as 100%. Preliminary experiments showed that the mean difference was 30% between the duplicates of the same specimen and 19% between the percentage found at 24 hours and that calculated from the values obtained at 12 and 36 hours. Preliminary experiments on artificial mixtures of normal and mutant hemizygous bloods, incubated with 10-4 mg./ml. of primaquine or with 1 × 10^-4 M MSB, also showed good agreement at the 24-hour survival rate with the expected value based on the proportions of the two kinds of cells mixed. It was, therefore, assumed that the radioactive salt entered equally into normal and mutant erythrocytes and was equally eluted from both, and that the decay of radioactivity during the short time considered was caused chiefly by destruction of erythrocytes altered by the exposure to MSB.

As different quantities of the substance added to the same blood led to different amounts of destroyed erythrocytes, it was further assumed that there was a continuous variability in their resistance against MSB damage and that this variability might be reflected by the differences between the radioactivity percentage figures of the 24-hour samples obtained after incubation with subsequent MSB dilutions. Since we found good agreement between the values obtained from the blood of the donor tested against the same dilutions at different times in different recipients, we also postulated that the individual differences among the recipients would not introduce any serious bias.

Results
The results listed in Table II and summarized in Fig. 1 were essentially as follows: (1) Both in the normal and in the mutant hemizygous blood the resistance of the erythrocytes against MSB-damage behaves as a nearly normally distributed function of the logarithm of the MSB-concentration.

(2) There is a considerable difference between the ranges of concentration that damage the normal and those that damage the mutant erythrocytes, the latter being definitely lower.

(3) There is almost no overlapping of the two

TABLE II
STUDIES WITH RADIOACTIVE 14Cr*

<table>
<thead>
<tr>
<th>MSB 10^-4 M</th>
<th>Subject</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>≈</td>
<td>10^-5</td>
</tr>
<tr>
<td>1</td>
<td>90-9</td>
</tr>
<tr>
<td>1/2</td>
<td>92-3</td>
</tr>
<tr>
<td>1/4</td>
<td>94-3</td>
</tr>
<tr>
<td>1/8</td>
<td>96-8</td>
</tr>
<tr>
<td>x</td>
<td>98-8</td>
</tr>
<tr>
<td>10^{-2}</td>
<td>75-7</td>
</tr>
<tr>
<td>10^{-3}</td>
<td>52-5</td>
</tr>
<tr>
<td>10^{-4}</td>
<td>30-3</td>
</tr>
<tr>
<td>10^{-5}</td>
<td>18-2</td>
</tr>
<tr>
<td>10^{-6}</td>
<td>10-1</td>
</tr>
<tr>
<td>10^{-7}</td>
<td>8-0</td>
</tr>
<tr>
<td>10^{-8}</td>
<td>5-0</td>
</tr>
<tr>
<td>10^{-9}</td>
<td>0-0</td>
</tr>
</tbody>
</table>

* See text.
Erythrocytes in Heterozygotes for G6PD Deficiency

by the 24-hour radioactivity-decay at the MSB 1, varies little in three but significantly in one female. From the values entered in Table II it can be seen that the proportion of normal erythrocytes ranges from 24.6 to 57.0%, with a mean of 46.5%.

Discussion

We feel that our results suggest the existence of two distinct populations of erythrocytes in females heterozygous for strong Mediterranean G6PD-deficiency. One is more damaged by MSB than the other and behaves as the single one of strong G6PD-deficient males. The other on the contrary behaves exactly as the single population of G6PD-normal males.

As shown by Zacchello (1963), incubation with MSB gives rise to the appearance of methaemoglobin and of Heinz-bodies in normal erythrocytes, and, at a higher dosage, equal to that determining complete destruction in vivo, it causes GSH-disappearance and loss of potassium. Furthermore our heterozygote with the lowest G6PD-enzyme level has the highest proportion of erythrocytes damaged by the MSB 1 (Subject No. 6), lower only than those of the mutant hemizygotess. All this strongly suggests that the two populations also differ in their mean G6PD-activity, as first postulated by Beutler et al. (1962).

Though we have not tested homozygous females to show that they have a single population, we conclude that the results presented here do complete those of Beutler and Baluda (1964) and agree with the hypothesis of Lyon (1962) on the behaviour of X-linked genes.

Summary

Short-term in vivo survival studies were carried out with erythrocytes damaged in vitro by increasing doses of menadione sodium bisulphite from two normal and two mutant hemizygotess and from four heterozygous for strong Mediterranean glucose-6-phosphate dehydrogenase deficiency.

The rate of the erythrocyte destruction referred to the menadione salt concentration showed a unimodal distribution in the hemizygotess, in ranges considerably higher in the normal than in the mutant ones. In the heterozygotes it showed a bimodal distribution, with two distinct peaks corresponding to the modes of the former.

This is considered proof of the existence of a true erythrocyte mosaic in females heterozygous for the enzyme deficiency, as hypothesized by Lyon.
Sartori, Panizon, and Zacchello

REFERENCES


Bimodal distribution of erythrocytes in heterozygotes for strong Mediterranean glucose-6-phosphate dehydrogenase deficiency.

E Sartori, F Panizon and F Zacchello

doi: 10.1136/jmg.3.1.42